

us-10-038-854-37

GenCore version 5.1.6

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OM nucleic - nucleic search, using sw model

Run on: August 2, 2004, 14:00:47 ; Search time 32049 Seconds
(without alignments)
11691.489 Million cell updates/sec

Title: US-10-038-854-37

Perfect score: 8645

Sequence: 1 ttggcctcgggccagaatt.....actgtattactaacttta 8645

Scoring table: IDENTITY_NUC
Gapop 10.0 , Gapext 1.0

Searched: 3470272 seqs, 21671516995 residues

Total number of hits satisfying chosen parameters: 6940544

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : GenEmbl:*

1: gb_ba:*
2: gb_htg:*
3: gb_in:*
4: gb_om:*
5: gb_ov:*
6: gb_pat:*
7: gb_ph:*
8: gb_pl:*
9: gb_pr:*
10: gb_ro:*

11: gb_sts:*
12: gb_sy:*
13: gb_un:*
14: gb_vi:*
15: em_ba:*
16: em_fun:*
17: em_hum:*
18: em_in:*
19: em_mu:*
20: em_om:*
21: em_or:*
22: em_ov:*
23: em_pat:*
24: em_ph:*
25: em_pl:*
26: em_ro:*
27: em_sts:*
28: em_un:*
29: em_vi:*
30: em_htg_hum:*
31: em_htg_inv:*
32: em_htg_other:*
33: em_htg_mus:*
34: em_htg_pln:*
35: em_htg_rod:*
36: em_htg_mam:*
37: em_htg_vrt:*
38: em_sy:*
39: em_htgo_hum:*
40: em_htgo_mus:*
41: em_htgo_other:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	%			Description
	Query	Match	Length	
	Score	DB	ID	

1	8645	100.0	8645	6	AX662355	AX662355 Sequence
2	8616.6	99.7	8675	6	AX662353	AX662353 Sequence
3	8000.6	92.5	8473	6	AX662357	AX662357 Sequence
4	7895.8	91.3	8487	6	AX662359	AX662359 Sequence
5	6789.2	78.5	8964	10	AB025412	AB025412 Mus muscu
6	5749.4	66.5	7816	10	AF195418	AF195418 Mus muscu
7	5305.8	61.4	5309	9	AB040888	AB040888 Homo sapi
8	4561.4	52.8	5804	10	AK122513	AK122513 Mus muscu
9	4273.8	49.4	8816	5	AB026979	AB026979 Danio rer
10	3714.8	43.0	8575	6	AX921803	AX921803 Sequence
11	3614	41.8	3614	6	AX876525	AX876525 Sequence
12	3614	41.8	3614	6	BD156175	BD156175 Primer fo
13	3614	41.8	3614	9	AK001336	AK001336 Homo sapi
14	3589.8	41.5	8438	6	AX675551	AX675551 Sequence
15	3566.6	41.3	8409	5	GGA279031	AJ279031 Gallus ga
16	3566.6	41.3	8409	6	AX250068	AX250068 Sequence
17	3545.6	41.0	8797	6	AX250063	AX250063 Sequence
18	3545.6	41.0	8797	6	AX250066	AX250066 Sequence
19	3545.6	41.0	8797	10	AB025411	AB025411 Mus muscu
20	3518.4	40.7	8689	6	AX250067	AX250067 Sequence
21	3518.4	40.7	8689	10	AF086607	AF086607 Rattus no
22	3502.2	40.5	8354	6	AX556500	AX556500 Sequence
23	3482.2	40.3	9826	6	AX250008	AX250008 Sequence
24	3453.8	40.0	9729	6	AX250013	AX250013 Sequence
25	3451.6	39.9	8585	10	AB025413	AB025413 Mus muscu
26	3439.2	39.8	8645	6	AX600210	AX600210 Sequence
27	3270	37.8	3270	6	AX877449	AX877449 Sequence
28	3270	37.8	3270	6	BD156663	BD156663 Primer fo
29	3270	37.8	3270	9	AK001748	AK001748 Homo sapi
30	3267.6	37.8	9722	10	AF059485	AF059485 Mus muscu
31	3235.6	37.4	3394	9	AK125869	AK125869 Homo sapi
32	3158	36.5	7781	9	AB032953	AB032953 Homo sapi
33	3066.4	35.5	9264	5	AB026980	AB026980 Danio rer
34	3012.6	34.8	8118	5	GGA238613	AJ238613 Gallus ga
35	2991.8	34.6	8373	10	AB025410	AB025410 Mus muscu
36	2970.8	34.4	7713	9	HSM808325	BX648178 Homo sapi
37	2854.8	33.0	8297	9	AF100772	AF100772 Homo sapi
38	2599	30.1	6560	6	AX250065	AX250065 Sequence
39	2155.4	24.9	2157	6	AX876360	AX876360 Sequence

40	2155.4	24.9	2157	6	BD156088
41	2155.4	24.9	2157	9	AK027473
42	1978.8	22.9	8993	9	HSM806812
43	1671.2	19.3	8624	9	AB037723
44	1658.8	19.2	5583	10	AK122490
45	1614.8	18.7	184032	9	AC079226

BD156088 Primer fo
AK027473 Homo sapi
BX640737 Homo sapi
AB037723 Homo sapi
AK122490 Mus muscu
AC079226 Homo sapi

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Search completed: August 3, 2004, 06:21:49
Job time : 32211 secs
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us-10-038-854-37

GenCore version 5.1.6

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OM nucleic - nucleic search, using sw model

Run on: August 2, 2004, 20:38:12 ; Search time 3396 Seconds
(without alignments)
12481.639 Million cell updates/sec

Title: US-10-038-854-37

Perfect score: 8645

Sequence: 1 ttggcctcgggccagaatt.....actgtatttaactta 8645

Scoring table: IDENTITY_NUC
Gapop 10.0 , Gapext 1.0

Searched: 3222919 seqs, 2451570024 residues

Total number of hits satisfying chosen parameters: 6445838

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Published_Applications_NA:*

- 1: /cgn2_6/ptodata/2/pubpna/US07_PUBCOMB.seq:*
- 2: /cgn2_6/ptodata/2/pubpna/PCT_NEW_PUB.seq:*
- 3: /cgn2_6/ptodata/2/pubpna/US06_NEW_PUB.seq:*
- 4: /cgn2_6/ptodata/2/pubpna/US06_PUBCOMB.seq:*
- 5: /cgn2_6/ptodata/2/pubpna/US07_NEW_PUB.seq:*
- 6: /cgn2_6/ptodata/2/pubpna/PCTUS_PUBCOMB.seq:*
- 7: /cgn2_6/ptodata/2/pubpna/US08_NEW_PUB.seq:*
- 8: /cgn2_6/ptodata/2/pubpna/US08_PUBCOMB.seq:*
- 9: /cgn2_6/ptodata/2/pubpna/US09A_PUBCOMB.seq:*
- 10: /cgn2_6/ptodata/2/pubpna/US09B_PUBCOMB.seq:*

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11: /cgn2_6/ptodata/2/pubpna/US09C_PUBCOMB.seq:*
12: /cgn2_6/ptodata/2/pubpna/US09_NEW_PUB.seq:*
13: /cgn2_6/ptodata/2/pubpna/US09_NEW_PUB.seq2:*
14: /cgn2_6/ptodata/2/pubpna/US10A_PUBCOMB.seq:*
15: /cgn2_6/ptodata/2/pubpna/US10B_PUBCOMB.seq:*
16: /cgn2_6/ptodata/2/pubpna/US10C_PUBCOMB.seq:*
17: /cgn2_6/ptodata/2/pubpna/US10_NEW_PUB.seq:*
18: /cgn2_6/ptodata/2/pubpna/US60_NEW_PUB.seq:*
19: /cgn2_6/ptodata/2/pubpna/US60_PUBCOMB.seq:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match %	Length	DB	ID	Description
1	8645	100.0	8645	17	US-10-038-854-37	Sequence 37, Appl
2	8616.6	99.7	8675	17	US-10-038-854-35	Sequence 35, Appl
3	8000.6	92.5	8473	17	US-10-038-854-39	Sequence 39, Appl
4	7895.8	91.3	8487	17	US-10-038-854-41	Sequence 41, Appl
5	3714.8	43.0	8575	13	US-10-072-012-143	Sequence 143, App
6	3614	41.8	3614	13	US-10-342-887-1743	Sequence 1743, Ap
7	3614	41.8	3614	13	US-10-172-118-1743	Sequence 1743, Ap
8	3589.8	41.5	8438	13	US-10-042-865-1	Sequence 1, Appli
9	3566.6	41.3	8409	9	US-09-808-602-79	Sequence 79, Appl
10	3566.6	41.3	8409	10	US-09-800-198-67	Sequence 67, Appl
11	3545.6	41.0	8797	9	US-09-808-602-74	Sequence 74, Appl
12	3545.6	41.0	8797	9	US-09-808-602-77	Sequence 77, Appl
13	3545.6	41.0	8797	10	US-09-800-198-62	Sequence 62, Appl
14	3545.6	41.0	8797	10	US-09-800-198-65	Sequence 65, Appl
15	3518.4	40.7	8689	9	US-09-808-602-78	Sequence 78, Appl
16	3518.4	40.7	8689	10	US-09-800-198-66	Sequence 66, Appl
17	3502.2	40.5	8354	13	US-10-383-201-43	Sequence 43, Appl
18	3502.2	40.5	8354	13	US-10-029-020-13	Sequence 13, Appl
19	3482.2	40.3	9826	9	US-09-808-602-7	Sequence 7, Appli
20	3482.2	40.3	9826	10	US-09-800-198-7	Sequence 7, Appli
21	3466.4	40.1	9695	16	US-10-144-194A-81	Sequence 81, Appl

22	3453.8	40.0	9729	9	US-09-808-602-12
23	3453.8	40.0	9729	10	US-09-800-198-12
24	3264.8	37.8	9058	16	US-10-144-194A-79
25	3249.6	37.6	8355	13	US-10-383-201-55
26	2854.8	33.0	12880	16	US-10-295-027-927
27	2599	30.1	6560	9	US-09-808-602-76
28	2599	30.1	6560	10	US-09-800-198-64
29	912.8	10.6	3217	15	US-10-198-846-13976
30	766.4	8.9	791	13	US-09-823-245A-85
31	706	8.2	2496	9	US-09-808-602-75
32	706	8.2	2496	10	US-09-800-198-63
33	663.8	7.7	1399	13	US-10-383-201-49
34	663.8	7.7	1399	13	US-10-383-201-59
35	622	7.2	1392	13	US-10-383-201-45
36	622	7.2	1392	13	US-10-383-201-53
37	621.2	7.2	1476	13	US-10-383-201-41
38	617.8	7.1	1371	13	US-10-383-201-51
39	584.4	6.8	1755	10	US-09-998-966-17
40	584.4	6.8	1755	15	US-10-004-415-17
41	584.4	6.8	1755	16	US-10-384-974-17
42	556.6	6.4	1973	9	US-09-864-761-4526
43	423	4.9	829	13	US-10-383-201-47
44	423	4.9	829	13	US-10-383-201-61
45	421.6	4.9	452	9	US-09-563-817-960

Sequence 12, App1
Sequence 12, App1
Sequence 79, App1
Sequence 55, App1
Sequence 927, App
Sequence 76, App1
Sequence 64, App1
Sequence 13976, A
Sequence 85, App1
Sequence 75, App1
Sequence 63, App1
Sequence 49, App1
Sequence 59, App1
Sequence 45, App1
Sequence 53, App1
Sequence 41, App1
Sequence 51, App1
Sequence 17, App1
Sequence 17, App1
Sequence 17, App1
Sequence 4526, Ap
Sequence 47, App1
Sequence 61, App1
Sequence 960, App

Q

Q